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PATENT

CERTIFICATE OF UTILITY



No. 12236*01

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Intellectual Property Code – Book VI

REQUEST FOR GRANT

DATE OF SUBMISSION OF THE DOCUMENTS NATIONAL REGISTRATION No. PLACE OF FILING DATE OF FILING	Jacques VESIN L'AIR LIQUIDE S.A. 75, quai d'Orsay 75321 PARIS CEDEX 07 France
Your file references: S6510JV	

1 NATURE OF THE APPLICATION							
Patent application							
2 TITLE OF THE INVENTION							
	Method for processing aluminum in a rotary or reverberatory furnace						
3 PRIORITY DECLARATION OR APPLICATION FOR THE BENEFIT OF THE FILING DATE OF A PRIOR FRENCH APPLICATION	<table border="1"> <thead> <tr> <th>Country or company</th> <th>Date</th> <th>No.</th> </tr> </thead> <tbody> <tr> <td colspan="3"> </td> </tr> </tbody> </table>	Country or company	Date	No.			
Country or company	Date	No.					
4-1 APPLICANT							
Name Handled by Street Postcode and town Country Nationality Legal form SIREN No. APE-NAF Code Telephone No. Fax No. Email	L'AIR LIQUIDE, SOCIÉTÉ ANONYME À DIRECTOIRE ET CONSEIL DE SURVEILLANCE POUR L'ETUDE ET L'EXPLOITATION DES PROCÉDÉS GEORGES CLAUDE Jacques VESIN 75, quai d'Orsay 75321 PARIS CEDEX 07 France France Société Anonyme 552 096 281 241A 01 40 62 57 32 01 40 62 56 95 jacques.vesin@airliquide.com						

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6 DOCUMENTS AND FILES ATTACHED		Electronic file	Pages	Details
Patent text		textebrevet.pdf	11	D 8, C 2, AB 1
Drawings		dessins.pdf	4	page 4, figures 3, Abstract: page 4, Fig.2
Designation of the Inventor				
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Method of payment		Debit to client account No.		
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8 SEARCH REPORT				
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062 Electronic filing		EURO	0.00	1.00
063 Search report		EURO	320.00	1.00
068 Claim from the 11th		EURO	15.00	2.00
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Function

L'AIR LIQUIDE, SOCIÉTÉ ANONYME À DIRECTOIRE ET CONSEIL DE SURVEILLANCE POUR L'ETUDE ET L'EXPLOITATION DES PROCÉDÉS GEORGES CLAUDE (Applicant 1)



PATENT CERTIFICATE OF UTILITY

Electronic receipt of the submission

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APPLICANT

Name or company name	L'AIR LIQUIDE, SOCIÉTÉ ANONYME À DIRECTOIRE ET CONSEIL DE SURVEILLANCE POUR L'ETUDE ET L'EXPLOITATION DES PROCÉDÉS GEORGES CLAUDE
Number of applicant(s)	1
Country	FR

TITLE OF THE INVENTION

Method for processing aluminum in a rotary or reverberatory furnace

DOCUMENTS SENT

package-data.xml Design.PDF FR-office-specific-info.xml dessins.pdf	Requetefr.PDF ValidLog.PDF application-body.xml indication-bio-deposit.xml	fee-sheet.xml textebrevet.pdf request.xml
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Method for processing aluminum in a rotary or
reverberatory furnace

5 The present invention relates to a method for
processing aluminum in a furnace, in which an aluminum-
containing material and, optionally, one or more salts
and/or slag and/or recycled dross, are introduced into
the furnace, this material is melted by heating using
10 at least one burner supplied with oxidizer and with
fuel, in order to obtain molten aluminum possibly
covered with a slag comprising, in particular, alumina,
and the carbon monoxide and/or hydrogen concentration
is measured in the furnace atmosphere or in the flue
15 gases.

In the field of secondary aluminum melting, this
operation is carried out in a rotary or reverberatory
furnace. Although this melting process may be
20 continuous, it is usually carried out in batches: the
materials are charged into the furnace, in one or more
successive cycles, before the molten metal is poured
toward its place of use. For this purpose, the molten
metal must have a temperature of about 740°C. Above
25 750°C, the rate of oxidation of the molten aluminum
increases considerably, and almost exponentially.

During a melting cycle, the initial period can
first be distinguished, when the materials are solid,
30 allowing the absorption of a large quantity of heat,
with the aluminum melting at about 660°C.

Irrespective of the type of furnace used, a slag
or "dross" is found on the melt surface. A distinction
35 is normally drawn between, on the one hand, what is
called "black" dross, which is a mixture of salts (if
the process uses salt), of various waste solid organic
compounds (polycyclic aromatic hydrocarbons, soot,

etc.), generated by the pyrolysis of the organic, and more generally, nonmetallic materials initially present in the charge, of aluminum oxide and aluminum imprisoned in the oxide, and, on the other hand what is called "white" dross, composed exclusively of aluminum oxides and aluminum, produced in the furnaces processing "clean" charges consisting exclusively of metal alloys intended for melting. It should be observed that the production of "black" dross is accompanied by the emission, in the flue gases, of large quantities of volatile organic compounds (VOC), rich in unburnt hydrocarbons.

This slag or dross contains a certain quantity of lost or oxidized metal also called "loss on ignition" which represents a non-negligible loss of material for the aluminum producer, and which should be reduced to the minimum in order to increase the profitability of the melting process. To reduce this oxidation, it is known to maintain the temperature of the aluminum melt to a value below about 750°C. However, this method remains empirical, because hot spots may appear on the surface, causing local oxidation.

Other known solutions prevent the oxidation by decreasing the contact of the metal surface with an oxidant.

Thus, document JP 58-227706 proposes to use the measurement of the CO and H₂ contents in the flue gases to ensure that, in a nonferrous metal smelter, the burners installed operate in substoichiometric mode in a range of values of the ratio of the oxidant to fuel flow rate ranging between 95% and 100%, part of the fuel remaining unconsumed.

Document EP 962 540 describes a combustion process for melting a metal in a furnace, in which an oxygen

rich gas is sent to the furnace, above the flame of a burner, in contact with the liquid metal.

5 The burner, operating in substoechiometric mode, produces a reducing flame that creates a barrier between the oxygen rich gas and the melt surface.

10 Document US 5 563 903 describes a method in which an inert or reducing gas creates a barrier between the aluminum melt surface and a combustion zone located in the upper part of the furnace.

15 Document US 3 759 702 describes a method in which the melting initially takes place in the open air, with a burner moving above the surface of the materials to be melted. The burner flame is slightly substoechiometric, hence reducing.

20 All these methods yield incomplete results and are applied throughout the duration of the melting, and not only when a risk of aluminum oxidation exists.

25 There is a need today for a method for processing aluminum that is applicable even if the first phase does not produce "black" dross and hence high VOC emissions, while limiting the oxidation of the aluminum during the final phase of the method.

30 The method of the invention serves to solve the problem stated and decrease the formation of aluminum oxides.

35 It is characterized in that the oxidizer supplied to at least one burner comprises over 10% by volume of oxygen, preferably over 21% by volume of oxygen, and in that the method comprises a final phase for decreasing the oxidation of the aluminum, during which the oxidizer flow rate is substantially constant while the

flow rate of fuel injected into at least one burner is a function of the carbon monoxide and/or hydrogen concentration in the atmosphere or the flue gases or vice versa, (that is, the fuel flow rate is substantially constant and the oxidizer flow rate depends on the CO and/or H₂ concentration comprised within the same limits as those defined below), this carbon monoxide and/or hydrogen concentration being between 3 vol% and 15 vol% (vice versa means the possibility in which the fuel flow rate is constant and the oxidizer flow rate depends on the CO and/or H₂ concentration).

Preferably, the oxidizer comprises over 88 vol% of O₂, preferably over 95 vol% of O₂. More preferably, the oxidizer is industrially pure oxygen.

The fuel may be any hydrocarbon or a light or heavy fuel oil (with an appropriate fuel oil spray system in the burner), and preferably natural gas, methane, propane, etc. is used. The volumetric ratio of oxygen to fuel is maintained between 1 and 5, preferably between 1.5 and 3.

According to one variant of the invention, the CO and/or H₂ concentration is maintained substantially constant throughout this oxidation limitation phase at a value of between 6 vol% and 10 vol% (the setpoint C2 during this final phase is therefore preferably set at a value within this range).

In general, the oxidation limitation phase is preceded by a VOC combustion phase during which substantially all the organic compounds present in the material are destroyed by pyrolysis, possibly (but not necessarily) followed by a stabilization phase.

Preferably, the VOC combustion phase terminates when the value of the ratio R of the volumetric flow rate of oxygen contained in the oxidizer to the volumetric flow rate of fuel during this phase falls below a threshold value S , defined below. In general, to contend with temporary fluctuations, the conditions of this hydrocarbon combustion phase are maintained for a further period Δt (between 5% and 20% of the hydrocarbon combustion phase in question) in order to confirm the passage of a value $R < S$, before entering the second phase by changing the setpoint (from a value $C1$ to a value $C2$) of the CO measured in the furnace or the flue gases, (considering the possible fluctuations of the variations in the CO concentration, as exemplified in Fig. 2, for example). This phase of duration Δt is referred to below as the detection phase.

According to a preferred embodiment of the invention, the method comprises two phases (which can be repeated several times before the liquid aluminum is poured) possibly separated by a generally short detection phase, intended to confirm the complete destruction of the organic compounds.

Preferably, during the first phase, the CO concentration in the furnace atmosphere and/or the flue gases, is between 0.1 vol% and 5 vol% (setpoint $C1$). The lower limit is in fact determined so that the flue gases (or the atmosphere) contain a maximum of 1000 ppm of oxygen.

In general, it has been found that a CO value of about 0.5 vol% is appropriate. During this first phase, the goal is to obtain the least possible CO in the atmosphere, that is, to adjust the setpoint $C1$ of the CO concentration at the lowest possible value in

the 0.1 vol% to 5 vol% interval, while preserving a nonoxidizing atmosphere in the furnace.

5 In the absence of control, the CO and/or H₂ concentration rises above the setpoint C1 selected above. Conversely, during the final phase, the H₂ and/or CO concentration (if the invention is not applied) is lower than the setpoint C2, and one of the goals of the invention is to increase this
10 concentration.

Thus in the initial phase, it is generally advantageous, using the regulation of the setpoint C1, to decrease this CO and/or H₂ concentration, while in
15 the final phase, the regulation of the setpoint C2 is used to increase the CO and/or H₂ concentration.

According to the invention, the transition from one phase to the other is based on the detection of a
20 durable variation of the ratio R ((volumetric oxygen flow rate)/(volumetric fuel flow rate) = R) going from a value above the threshold S to a value below the threshold S .

25 The threshold S of the ratio R is defined by prior tests on the furnace in which the invention is implemented, as follows:

- the setpoint C1 to which the CO concentration in the flue gases and/or the atmosphere is regulated
30 (using a variation of the ratio R) is set at the lowest possible value for the furnace tested, its charge (usual) and the CO detector used (generally a laser diode according to the invention). This value is often about 0.1 vol%. With the furnace having a charge as
35 normally processed, the CO value is adjusted to this very low setpoint: after a certain interval, the ratio of the volumetric oxygen/fuel flow rates is stabilized.

The ratio R at which this stabilization is obtained is the threshold S defined above.

Thus in the method of the invention, the CO and/or
5 H₂ is regulated during the first phase (combustion of volatile organic compounds VOCs) about a setpoint of between 0.1 vol% and 5 vol% (0.5 vol% is often satisfactory), and the ratio R defined above is measured. When R decreases to less than S, regulation
10 is generally continued to the setpoint C1, and then, after a few more minutes (after making sure that the phase has changed, that is, that all the VOCs are burnt), the setpoint is changed to a new setpoint C2 of between 3 vol% and 15 vol%, preferably between 6 vol%
15 and 10 vol%, the beginning of the final phase, in which the ratio R remains below S until the molten metal is poured.

On the contrary, after a certain period in the
20 final phase, aluminum waste (cans, etc.) can be reintroduced in order to regenerate an initial phase in which regulation again proceeds about the setpoint C1, and then, as previously, about the setpoint C2.

25 According to a variant of the invention only comprising one final phase (clean charge without VOC emission), regulation is directly carried out about a setpoint C2 of between 3 vol% and 15 vol% of CO, a value higher than the CO concentration in the same
30 furnace with the same charge, in the absence of regulation to a setpoint by CO and/or H₂.

The aluminum oxidation limitation phase terminates with the reintroduction of a new charge of aluminum-
35 containing material in the furnace, or with the pouring of the liquid aluminum toward its point of use.

In the context of the invention, the aluminum-containing material may be in particular, for example, aluminum in ingots, chips from the turning of aluminum parts, beverage and food cans, scrap, production waste, dross, aluminum-containing slag, and in general, any aluminum-containing material. Obviously, the invention also applies to liquid aluminum temperature-holding furnaces.

10 The invention will be better understood from the following embodiments, provided as nonlimiting examples, jointly with the figures which show:

15 - Figure 1, a schematic view of a furnace with a single burner shown, the flue gas analysis and burner control;

 - Figure 2, a diagram explaining the Phases I and II of a method of the invention;

20 - Figure 3, a graph explaining the variations in (CO) as a function of time showing the phases (or sub-phases) of the method of the invention.

 Figure 1 is a schematic view of a furnace (cross section) and of the control system according to the invention.

25

 A burner 10 creates a flame 2 which heats and melts the metal 3, in liquid form. The flue gases 4 issuing from the furnace 1 and produced by the combustion, particularly by the burner, are removed via the line 18, in which the CO and/or H₂ detectors 5 and 6 (known per se) are respectively placed, for measuring the CO and/or H₂ concentration in said flue gases. The signal issuing from each of the detectors 5 and 6 is transmitted via the connecting line to a control unit 8 of which the operation is explained below. The burner 10 is supplied respectively with oxidizer 13 and fuel 14 via controlled valves 12 and 11 respectively (mass flowmeters, for example) for delivering an appropriate

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flow rate of oxidizer and fuel to the burner. This flow rate is controlled by the control device 8, via the connecting line 15. The connecting lines 17 and 16 transmit the measurement of the opening of the valves 12 and 11 to the control system 8, which also receives data on the temperature of the molten metal 3 via a sensor. The control system 8 comprises an adjustment of the CO (and/or H₂) concentration setpoint, which can be modified as a function of time, particularly to go from the value C1 to the value C2.

According to whether the measurement of the CO and/or H₂ concentration transmitted by the sensors 5 and/or 6 to the control device 8 is higher or lower than said setpoint, it generates a control signal via the connection 15 to the controlled valves 12 and 11 which adjust the injection of oxidizer 13 and fuel 14 to obtain a decrease or increase in the carbon monoxide and/or hydrogen concentration in the flue gases.

20

Figure 2 shows the typical variations of the ratio R (the flow rate of oxidizer or, preferably, the flow rate of fuel, is kept constant) and of the CO and/or H₂ concentration in the flue gases, in the context of the two-phase management of the melting cycle described above. During phase I, the CO and/or H₂ setpoint is set at the value of C1 of between 0.1 vol% and 5 vol%, whereas in phase II, this setpoint is adjusted to the value C2, substantially high, between 3 vol% and 15 vol%, with, in all cases, in the same successive cycle of phases I and II, C1 = C2 and C1 < C2 (if C1 = C2, there would only be a single phase). As the melting begins, in order to servocontrol the CO and/or H₂ to the setpoint, the controller increases the value of the ratio R in order to supply excess oxidizer to burn the VOCs. The production of VOC by the charge and their combustion reaches a peak and then drops to zero at the end of phase I. The ratio R follows this trend by

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rising to a peak, denoted R_{MAX} , and then decreasing in phase I. Because of the setpoint $C1$, when the VOCs are almost entirely consumed, the ratio R necessarily and definitively passes through the threshold S , previously defined, and then drops to a minimum, denoted R_{MIN} . When the ratio R reaches the threshold S , a small quantity of VOC actually remains in the charge, generating a level of CO and/or H_2 lower than the setpoint $C1$. From this time on, the controller adjusts the ratio R , decreasing it further (production of CO and/or H_2 by the burner) to compensate for the deviation from this setpoint $C1$. All the VOCs are burnt when the ratio reaches R_{MIN} . It is then time to change the control strategy because phase II begins. The invention consists in using the time $t1$ of crossing the threshold S coupled with a detection time Δt of between 5% and 20% of the melting phase I considered (which terminates at time $t1$) to engender a change in control strategy, at time $t2$ defined by $t2 = t1 + \Delta t$. At this time, the setpoint becomes equal to $C2$.

According to a variant of the invention, it is possible, conversely, to detect the introduction of fresh materials into the furnace and to switch from the oxidation limitation phase to the VOC combustion phase. In fact, considering a charge of aluminum during phase II, charge-related VOCs are released and the controller adjusts the ratio R by increasing it (decreased production of CO and/or H_2 by the burner), to compensate for the deviation from this setpoint $C2$. Similarly, the ratio R exceeds the threshold S at a given time $t3$, indicating that the physical process is in phase I, so that the change in control strategy, ordered by the indicator, takes place at time $t4$ defined by $t4 = t3 + \Delta t$. At this time $t4$, the setpoint is therefore restored to the value $C1$ defined above. For the process control unit, the end of phase II is hence time $t4$.

Embodiment

5 In a rotary furnace equipped with a 13 MW burner,
27 tonnes of salt and 27 tonnes of aluminum scrap are
charged, the charge is heated until 2550 Sm³ of natural
gas are consumed, then 65 tonnes of aluminum scrap are
again charged (beverage cans in this example) and the
charge is heated until an additional 850 Sm³ of natural
10 gas are consumed. 35 tonnes of aluminum scrap
(aluminum lathe turnings) are then again charged and
the charge heated until an additional 1350 Sm³ of
natural gas are consumed, 99 tonnes of aluminum are
then poured, the salts and dross present in the furnace
15 are then heated until a total of 5500 Sm³ of natural gas
are consumed. The aluminum remaining in the furnace is
then poured: 9 tonnes of aluminum.

Figure 3 shows the variations in CO measured in
20 the flue gases and the variations in the ratio R,
defined above, during the melting of the two aluminum
charges described above. In these curves, the effect
of the automatic change in strategy is clearly
observable. In fact, the indicator according to the
25 invention actuates the mid-fusion phase change at time
t₂ because the first charge consists of painted
beverage cans, hence with a high concentration of
organic compounds. Furthermore, during the melting of
the second charge, which is poorer in organic compounds
30 (aluminum lathe turnings), the indicator controls the
phase change at t'₂ shortly after the phase begins,
thereby optimizing the initiation of the control
strategy adapted to phase II. This indicator serves to
decrease the oxidation losses regardless of the type of
35 charge entering the furnace, as shown by the table
below:

1st filing

- 12 -

	Type of material charged	Unit	Method without control	Method according to the invention
Weight of oxidized aluminum	cans	kg	1800	1750
Weight of oxidized aluminum	turnings	kg	2000	1750

CLAIMS

1. A method for processing aluminum in a furnace,
5 in which an aluminum-containing material and, optionally, one or more salts, are introduced into the furnace, this material is melted by heating using at least one burner supplied with oxidizer and with fuel, in order to obtain molten aluminum possibly covered
10 with a slag comprising, in particular, alumina and at least one salt, and the carbon monoxide CO and/or hydrogen H₂ concentration is measured in the furnace atmosphere or in the flue gases, characterized in that the oxidizer supplied to at least one burner comprises
15 over 10% by volume of oxygen, preferably over 21% by volume of oxygen, and in that the method comprises a final phase for decreasing the oxidation of the molten aluminum, during which the oxidizer flow rate is substantially constant while the flow rate of fuel
20 injected into at least one burner is a function of the carbon monoxide and/or hydrogen concentration in the atmosphere or the flue gases or vice versa, this carbon monoxide and/or hydrogen concentration being regulated to a setpoint C2 of between 3 vol% and 15 vol%.

25

2. The method as claimed in claim 1, characterized in that the oxidizer comprises over 88 vol% of O₂, preferably over 95 vol% of O₂.

30 3. The method as claimed in either of claims 1 and 2, characterized in that the oxidizer is industrially pure oxygen.

4. The method as claimed in claim 1, characterized
35 in that the fuel is selected from natural gas, hydrocarbons, and light or heavy fuel oil.

5. The method as claimed in one of claims 1 to 4, characterized in that the volumetric ratio of oxygen to fuel is maintained between 1 and 5, preferably between 1.5 and 3.

5 6. The method as claimed in one of claims 1 to 5, characterized in that the carbon monoxide and/or hydrogen concentration is maintained substantially constant throughout this oxidation limitation phase at
10 a value C2 of between 3 vol% and 15 vol%, preferably between 6 vol% and 10 vol%.

7. The method as claimed in claims 1 to 6, characterized in that the oxidation limitation phase is
15 preceded by a hydrocarbon combustion phase during which substantially all the organic compounds present in the material are destroyed by pyrolysis.

8. The method as claimed in claim 7, characterized
20 in that the hydrocarbon combustion phase is considered to terminate when the measured value of the ratio R of the volumetric flow rate of oxygen to the volumetric flow rate of fuel falls below a predefined value S.

25 9. The method as claimed in claims 7 and 8, characterized in that a stabilization phase takes place with a measured CO and/or H₂ concentration regulated to the setpoint C1, this phase terminating when the ratio R reaches its minimum.

30 10. The method as claimed in one of claims 1 to 9, characterized in that the aluminum oxidation limitation phase terminates with the reintroduction, into the furnace, of a new charge of aluminum-containing
35 material.

11. The method as claimed in one of claims 1 to 10, characterized in that the CO concentration is measured using a laser diode.

- 5 12. The use of a laser diode for measuring CO in a method as claimed in at least one of the preceding claims.

1/3

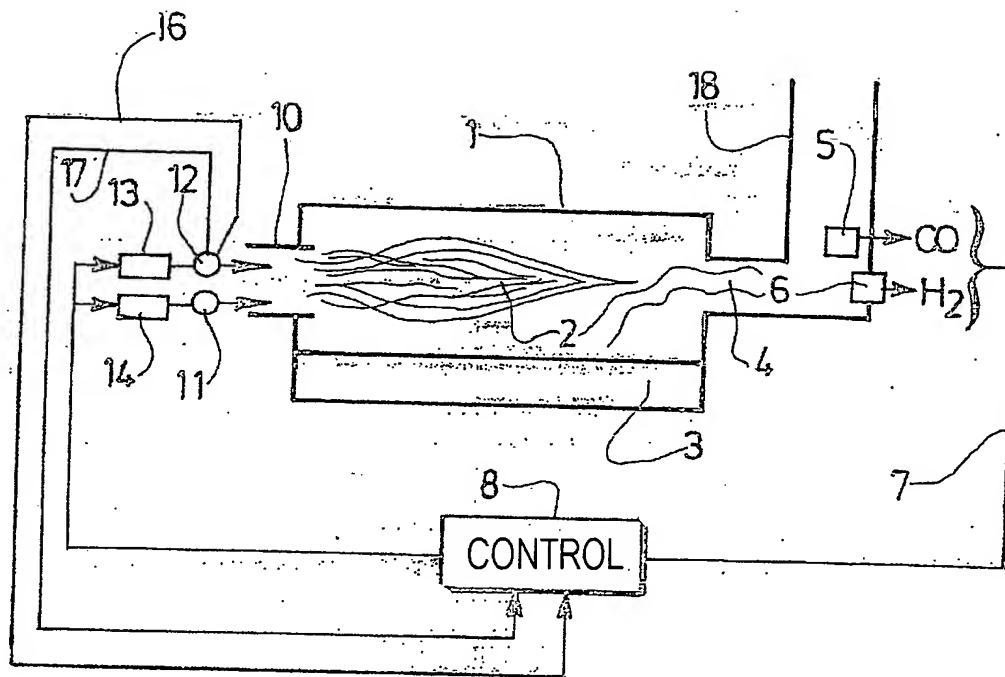


FIG.1

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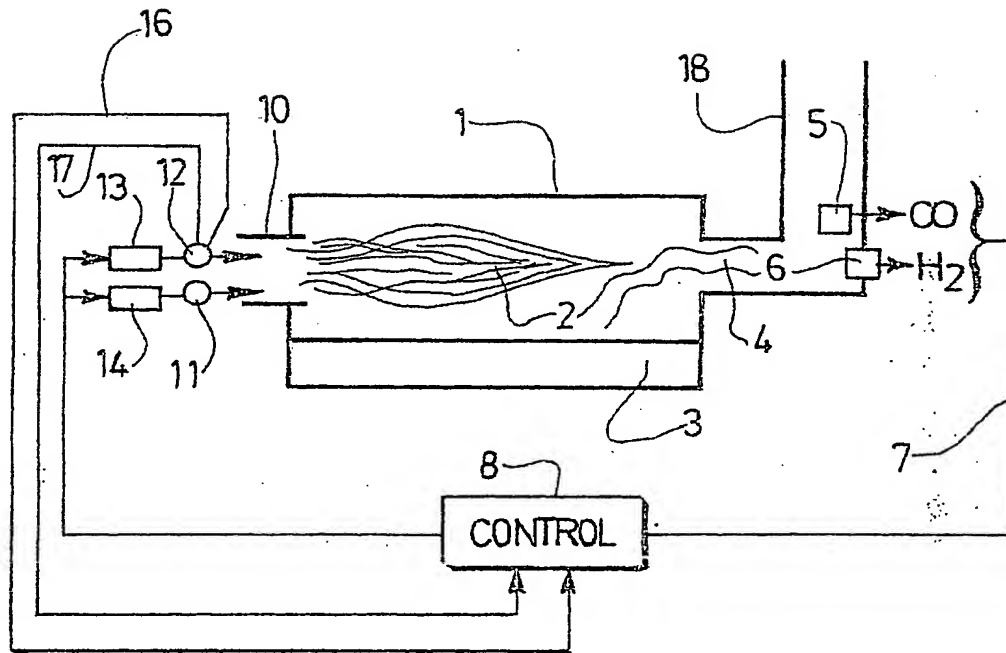


FIG.1

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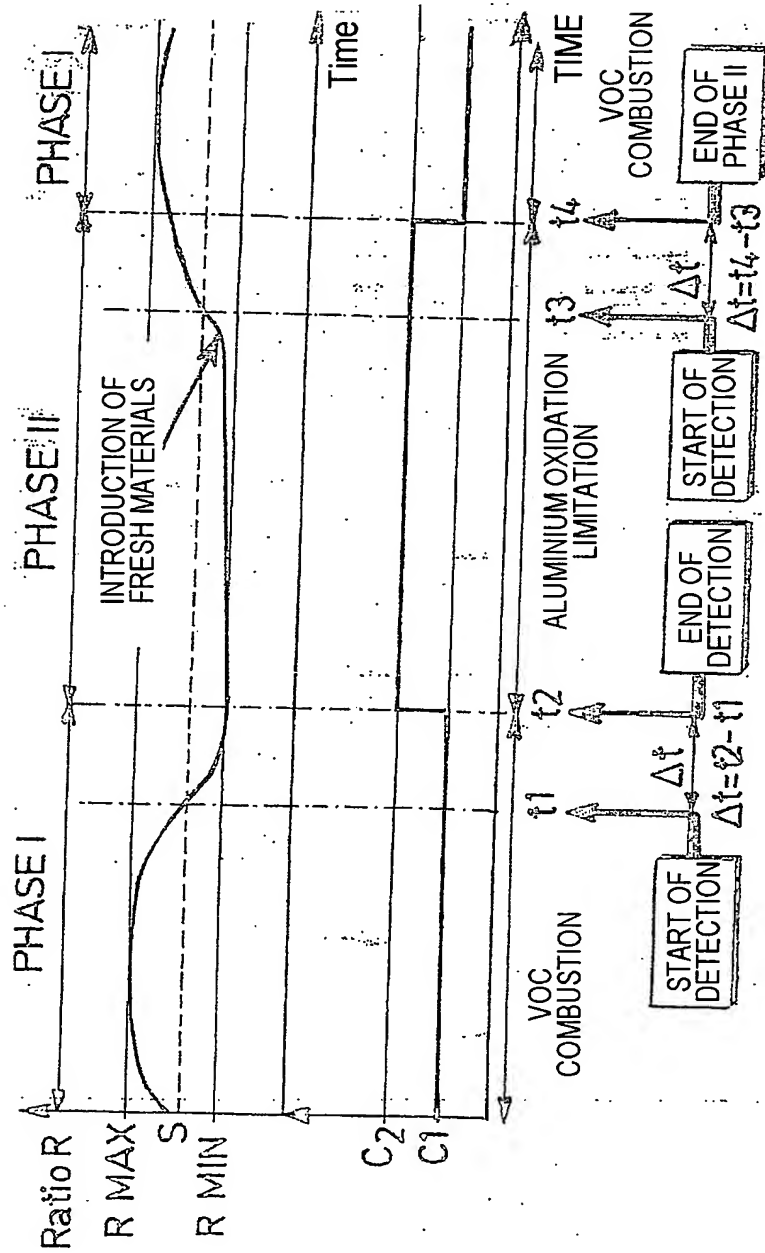


FIG. 2

2/3

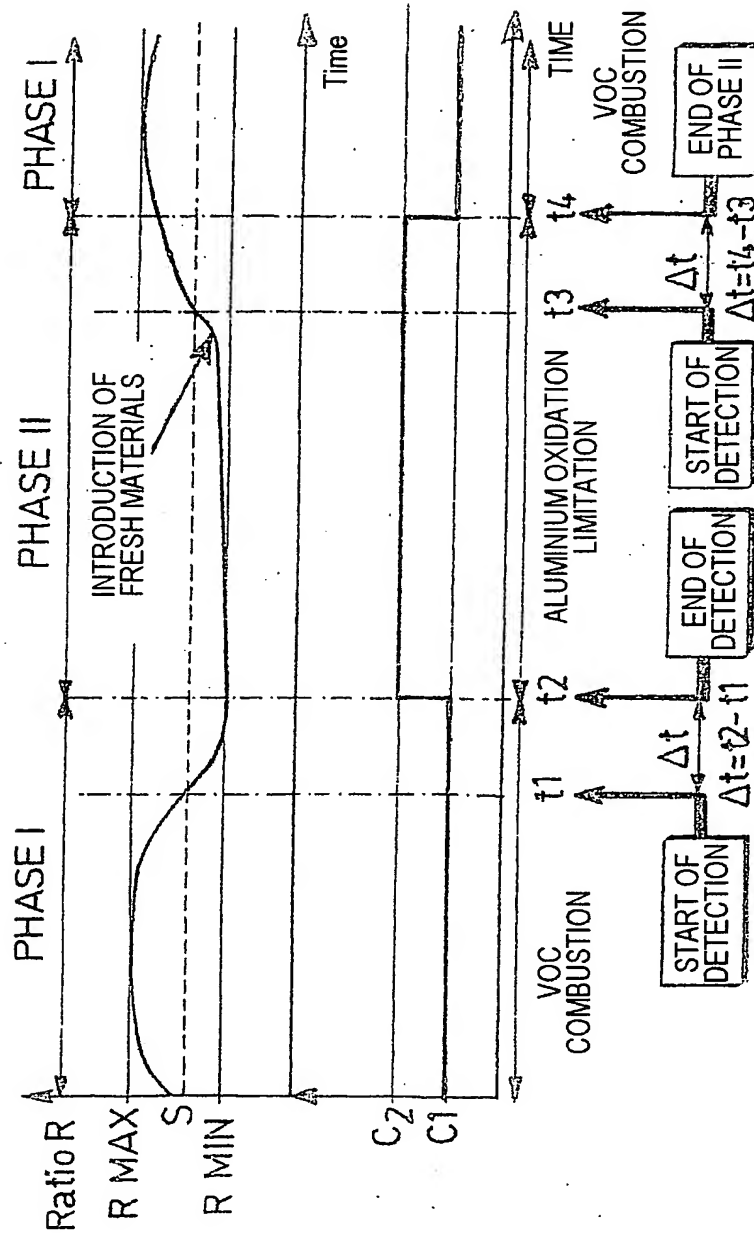
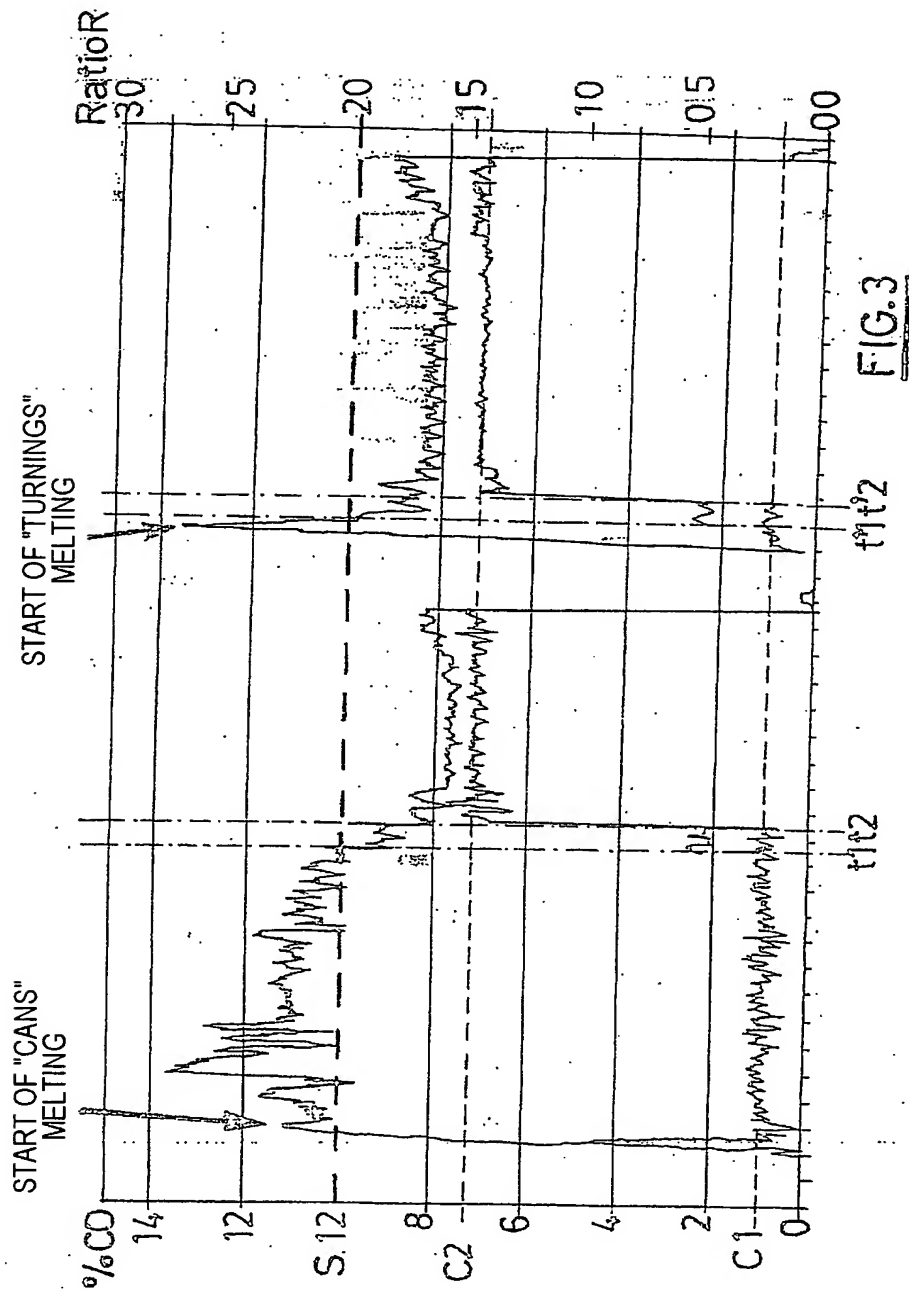


FIG.2

3/3



3/3

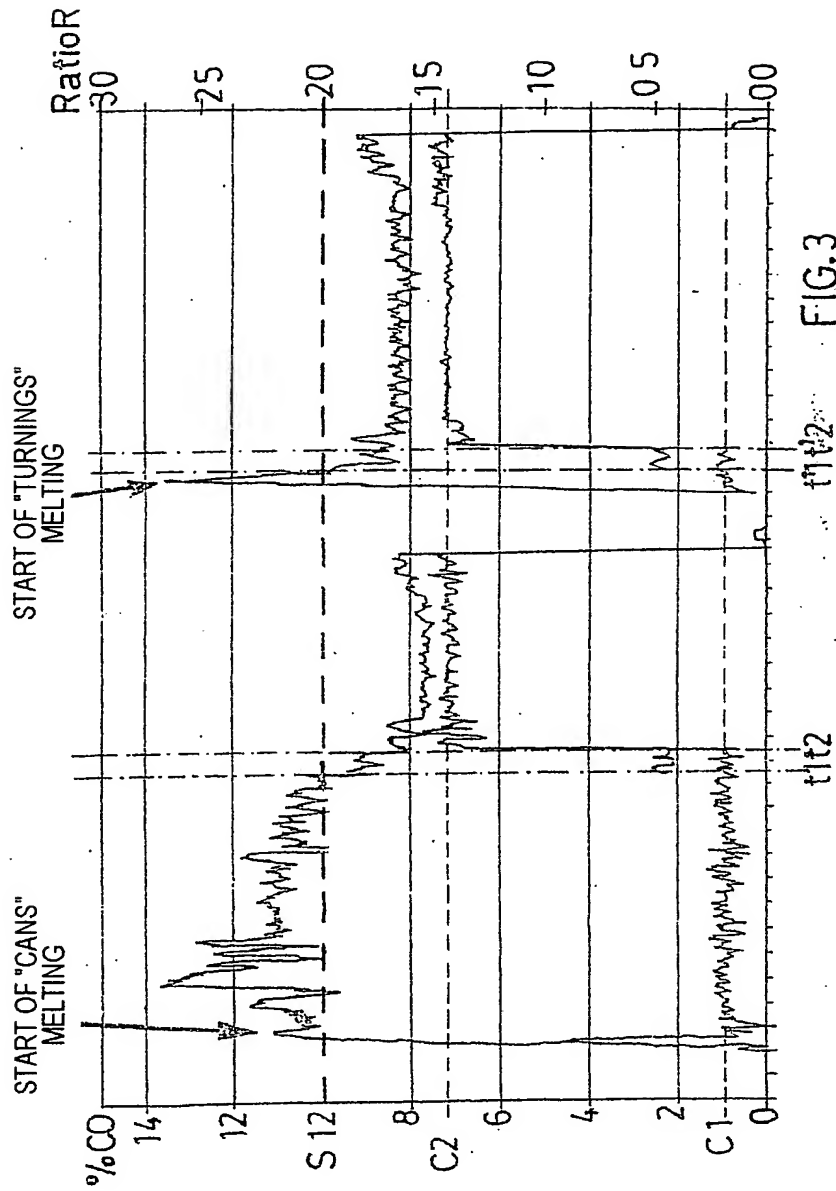


FIG.3



PATENT

CERTIFICATE OF UTILITY



No. 12236*01

DESIGNATION OF THE INVENTOR

Your file references	S6510JV
National Registration No.	
TITLE OF THE INVENTION	
	Method for processing aluminum in a rotary or reverberatory furnace
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Function

L'AIR LIQUIDE, SOCIÉTÉ ANONYME À DIRECTOIRE ET CONSEIL DE SURVEILLANCE POUR L'ETUDE ET L'EXPLOITATION DES PROCÉDÉS GEORGES CLAUDE (Applicant 1)